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From: Einsmann, Juliet  
Sent: Tuesday, October 08, 2002 2:17 PM  
To: STIC-ILL  
Subject: please deliver (for 09/761579)

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Tripatara et al. Archives of biochemistry and biophysics (1999 Jul 1) 367(1)39-50

Lissens et al. Human Mutation (1996) 7(1) 46-51

Bonne et al. Pediatric research (1993 Mar) 33(3)284-8

Dahl et al. Human Genetics (1991 May) 87(1) 49-53

Huh et al. Journal of Biological Chemistry (1990 Aug 5) 265 (22) 13320-6

Kitano et al. Journal of inherited metabolic disease (1989) 12(2)91-107.

Juliet Einsmann  
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703 306 5824  
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Teach that  $E_{1\alpha}$  subunit mutations  
are related to severity of  
 $E_{1\alpha}$  deficiency in patients with  
PDH complex deficiency.  
Are silent, however as to  
the molecular basis of the  
mutation.

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Teach a series of mutations that  
cause changes in coding sequence of  
PDH E<sub>1</sub> gene (table 2) in  
patients w/ PDH complex deficiency

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Absence of a complex in

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